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LEUCINE-RICH REPEATS (LRRs) were first discovered in leucine-rich a2-glycoprotein, a protein of unknown function from human serum1. LRR-containing proteins now represent a diverse group of molecules with differing functions and cellular locations in a variety of organisms (Table I, Fig. 1). LRRs are usually present in tandem, and the number of LRR motifs ranges from one, in platelet glycoprotein IbB, to 30, in chaoptin. The most common length of an LRR is 24 residues, but repeats containing any number between 20 and 29 residues are found. The recent X-ray structure analysis of ribonuclease inhibitor (RI) from pig liver2 (Fig. 2), a protein built entirely from LRRs, provided the first picture of the threedimensional architecture of LRRs.

Consensus sequences of LRRs and evolutionary implications

As the name implies, LRRs are distinguished by a consensus sequence consisting predominantly of leucines (Table 1). LRRs are easily overlooked by the commonly used methods for sequence comparison because they contain gaps, vary in length and amino acid composition, and therefore have a degenerate consensus sequence. Consequently, LRRs in many proteins were not detected for some time.

The consensus sequence compiled from all known LRR-containing proteins contains leucines or other aliphatic residues at positions 2, 5, 7, 12, 16, 21 and 24, and asparagine, cysteine or threonine at position 10 (Table I). Most proteins contain exclusively asparagine at position 10, but three have exclusively cysteine in this position (Table I). The functional and evolutionary significance of the variations of consensus residues at position 10 is not known.

Consensus sequences derived from LRRs in individual proteins often contain additional conserved residues in positions other than those mentioned above: mostly these are aliphatic and aromatic amino acids, sometimes glycines and prolines, but seldom other amino acids. The hydrophobic consensus residues in the carboxy-terminal parts of the repeats

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The leucine-rich repeat: a versatile binding motif

Bostjan Kobe and Johann Deisenhofer

Leucine-rich repeats are short sequence motifs present in a number of proteins with diverse functions and cellular locations. All proteins containing these repeats are thought to be involved in protein-protein interactions. The crystal structure of ribonuclease inhibitor protein has revealed that leucine-rich repeats correspond to β - α structural units. These units are arranged so that they form a parallel β-sheet with one surface exposed to solvent, so that the protein acquires an unusual, nonglobular shape. These two features may be responsible for the proteinbinding functions of proteins containing leucine-rich repeats.

are commonly spaced by 3, 4 or 7 residues.

About half of the genes encoding the known LRR-containing proteins have been sequenced. Gonadotrophin receptor genes contain several similarly sized exons in their LRR domains that are homologous in their exon-intron junction sequences³, suggesting that the LRR domain evolved by exon duplication and shuffling from a single prototypic exon corresponding to one LRR. Such an evolutionary relationship is less obvious in genes encoding other LRRcontaining proteins, but the introns often localize at similar positions in the repeats (most frequently between positions 1 and 6, less frequently between positions 11 and 18, and rarely elsewhere). The most likely mechanism of evolution of LRR-containing proteins would therefore comprise unequal crossovers and duplications of gene fragments corresponding to prototypic leucine-rich building blocks. Two evolutionary scenarios are possible: (1) an ancestral LRR module evolved first, and duplicated to give rise to individual members of the superfamily that later developed their own unique consensus sequences; or (2) the duplications occurred separately in each protein family and, owing to structural constraints, gave rise to similar, but unique, consensus sequences. These two schemes are not mutually exclusive and there is good reason to believe that the divergent and convergent scenarios have worked in conjunction to yield the LRR superfamily that exists today. The variation in the length of LRRs among proteins and the

presence of consensus sequences unique to a particular protein argue for a rich and complex evolutionary history that may have sprung from more than one root.

Three-dimensional structure of LRRs

The recently determined crystal structure of porcine ribonuclease inhibitor protein (RI)2 (Fig. 2a) has allowed us to examine the threedimensional architecture of LRRs. RI is a cytoplasmic protein that inhibits ribonucleases from the pancreatic superfamily by binding very tightly to an extensive surface area of ribonuclease containing its catalytic site4. The sequence of RI consists of 15 alternating LRRs of 28 (A-type) or 29 (B-type) residues each (Fig. 2b). Ninety percent of the structure is built of these repeat units. The short amino- and carboxyterminal segments display a weaker homology to the internal repeat motif.

In the RI molecule, LRRs correspond to β - α structural units, consisting of a short β-strand and an α-helix approximately parallel to each other (Fig. 2). All repeats, including the terminal segments, adopt very similar structures: All structural repeats contain 28 or 29 residues, except the amino-terminal repeat, which contains 25 residues. The structural units are arranged so that all the B-strands and the helices are parallel to a common axis, resulting in a nonglobular, horseshoe-shaped molecule with a curved parallel β-sheet lining the inner circumference of the horseshoe, and the helices flanking its outer circumference.

RN: Leu RN: U2 Big Dec Fibi Lun Pro Ost Plat Plat Yop Ipat

Protein (species)*	Function-ligand	Locationsb	Repeats	Length	Consensus sequence	PIR ^r entry
Part					5 10 15 20 25	
RNase inhibitor (porcine)	RNase inhibitor-RNase	Cytoplasm	15	28 (A)	.LE.L.LCLTCLaL	A31857
		•	•	29 (B)	.L.EL.LNLGD.GaLLP	02001
Leucine-rich a2-GP (human)	?-?	Serum	8	24	.LL.LNLLLL	NBHUA2
RNA1 (Saccharomyces cerevisiae)	RNA processing-?	Cytoplasm	8	29		
U2 anRNP A' (human)	Splicing-U2 snRNP	Nucleus	4	24	.LL.LNaaaa.	BVBYN1
Biglycan (human)	ECM binding-laminin.	ECM	8	24	.LL.aNaL	S03616
28,021 (121131)	fibronectin, TGF-8	ECIVI	•	24	.LL.LNIaa	A40757
Decorin (human)	ECM binding-collagen, fibronectin, thrombospondin, TGF-β	ECM	10	24	.LL.LNIVa	NBHUC8
Fibromodulin (bovine)	ECM binding-collagen.	ECM	11	24	.L. L.LNaaa	COEDOO
	fibronectin			24		\$05390
Lumican (chicken)	Comeal transparency-?	ECM	12	24	.LL.LNLa	A41748
Proteoglycan-Lb (chicken)	?-?	ECM	6	24	.L. a.LNIa	
Osteoinductive factor (bovine)	Bone morphogenesis-BMP	ECM	6	24		A41781
Platelet GP Iba (human)	Cell adhesion-vWF, thrombin	PM (EC)	7	24	.L.a.LNaP	A35272
Platelet GP V (human)	Cell adhesion-GP IX, GP Ib				.LL.LNLLP.GLL	NBHUIA
YopM (Yersinia pestis)		PM (EC)	14	24	.L.,L.L.,N-,L.,-LP.,LF.,L,	-
	Virulence factor-thrombin	IC + EC	12	20	.LL.aNLLPLPP	A33950
IpaH7.8 (Shigella flexneri)	3-3	?	6	20	.LL.VNLLPLP.	A35149
IpaH4.5 (Shigelia flexneri)	25	?	8	20	.LL.aNL.,-LP-,LP.	S18248
Toll (<i>Drosophila</i>)	Embryo development-?	PM (EC)	19	24	.LL.L. NL F	A29943
Slit (Drosophila)	Axon development-?	EC	19	24	.LL.LNIFL	A36665
Connectin (Drosophila)	Synapse development-?	PM (EC)	7	24	.LLNL . N Ia aF L	
Chaoptin (Drosophila)	Photoreceptor-cell development-7		30	24		S28464
Flightless-I (Drosophila)	Embryo development-?	PM (EC)	16	23	.LL.LNaaFa	A29944
Oligodendrocyte myelin GP (human)	Myellnation-?	PM (EC)	8	24	.LL.LS.NLaPaL .LL.LSNNaaL	A34210
CD14 (human)	Cell-surface receptor-LPS-LPB	PM (EC)		27		
irk (human)				27	.aL.LN	TDHUM4
	Receptor protein kinase-NGF	PM (EC)	2 .	23	.LL.LS.NL	TVHUTT
TrkB (mouse)	Receptor protein kinase-BDNF, NT-3	PM (EC)	3	23	.LL.aT.NLTST	S06943
frkC (porcine)	Receptor protein kinase-NT-3	PM (EC)	3	23	.LR.aNLSQNLS	440000
TMK1. (Arabidopsis thaliana)	Receptor protein kinase-?	PM (EC)		23 ·		A40026
LH-CG receptor (rat)	Signal transduction-LH, CG	PM (EC)		25 25	.ba.LNG.aPa.SL	JQ1674
SH receptor (rat)	Signal transduction-FSH				.LL.aTaF	A41343
(SH receptor (dog)		PM (EC)		25	.LL.aS.TLPaa	A34548
	Signal transduction—TSH	PM (EC)		25	.aL.a.NNa.S-aa	A40077
Adenylate cyclase (Saccharomyces cerevisiae)	Signal transduction-RAS	PM (cytoplasm)	20	23	.LL.LNaaaL	OYBY
FLR (Trypanosoma brucel)	7-7	?	18	23	.LL.LSGCaaaL	A36359
RAD1 (Saccharomyces cerevisiae)	DNA repair=RAD10	Nucleus				
AD7 (Saccharomyces cerevisiae)	DNA repair-?	?			.a.LaDINLPaN	DOBYO1
ORT100 (Arabidopsis thallana)	Recombination—?	•	-		.LL.aCaaa.	A25226
		Chloroplast			.LLNLNL.G.IP.S-a.S	A46260
RR1 (Saccharomyces cerevisiae)	Signal transduction-?	Cytoplasm		26	.La.LC.NaTDaLL	A41529
CR4 (Saccharomyces cerevisiae)	Transcription-?	?	4	23	.LL.aNLTLP.E-a	531286
ds22 (Schizosaccharomyces pombe)	Mitosis-dis2, sds21	Nucleus	11	22	.L.,L.a.,NI.,-aENa,.L	A38439
34 ribosome-binding protein (rat)	RM membranes-ribosome	RM membrane (cytoplasm)	4 :	24	LLOLNLLPFL	-
arboxypeptidase N (human)	Stabilization-catalytic subunit	Plasma	12	24		
nternalin (Listeria monocytogenes)					L.L.L.NLLPaPL	A34901
		Cell wall			LL.L.N-QISDI.PLLT	A39930
niB (Listeria monocytogenes)	?-?	?	6 2	22 .	LL.LNL.DILL 5 10 15 20 25	C39930
RR superfamily					5 10 15 20 25 LL.LN±.a±a±±±±a±±a±±,±±	

"A single example Is given for proteins where sequences from several species are known. Proteins with less than two LRRs are not shown. Abbreviations: RNase, ribonuclease; GP, glycoprotein; snRNP, small nuclear ribonucleoprotein particle; ECM, extracellular matrix; PM, plasma membrane; EC, extracellular; TGF, transforming growth factor; IC, intracellular; BMP, bone-morphogenic protein; vWF, von Willebrand factor; LPS-LPB, complex of lipopolysaccharide and lipopolysaccharide-binding protein; NGF, nerve growth factor; BDNF, brain-derived neurotrophic factor; NT-3, neurotrophin-3; LH, lutrophin; CG, choriogonadotrophin; FSH, follitrophin; TSH, thyrotrophin; T-R, trypanosomal leucine-rich protein; RM membrane, rough microsomal membrane.

Cellular distribution of the protein. For membrane-associated proteins, the location of the LRR domain is indicated in parentheses.

Total number of repeats is the number of occurrences of the a.a.a..N/C/T sequence, where 'a' represents A, V, L, I, F, Y or M; repeats shorter than 18 residues and isolated single repeats were not counted. Only the counted repeats were used to determine the consensus sequence. Because other authors may have used different criteria, the numbers of repeats, the lengths and the consensus sequences may vary from those originally reported.

The most common length of all the repeats in the protein is given. For some proteins, there are significant deviations from this number. RNase inhibitor is aligned so that two types of repeats (A-type and B-type) alternate in the sequence.

"The numbers above the sequences indicate the position in the repeat in reference to the consensus of porcine RNase inhibitor. One-letter code is used for amino acids. An amino acid is included in the consensus if present at that position in more than half of the repeats; 'a' represents A, V, L, I, F, Y or M, and is included in the consensus if these amino acids are present at that position in more than 80% of the repeats. Symbols used: '.', any amino acid; '--', gap; '±', amino acid may or may not be present at this position.

Protein Identification Resource sequence data bank. References for the proteins without PIR entries: Platelet GV, Hickey, M. J. et al. (1993) Proc. Natl Acad. Sci. USA 90, 8327–8331; Rightless-I, Campbell, H. D. et al. (1993) Proc. Natl Acad. Sci. USA 90, 11386–11390; p34 ribosome-binding protein, Ohsuml, T. et al. (1993) Biochem. J. 294, 465–472.

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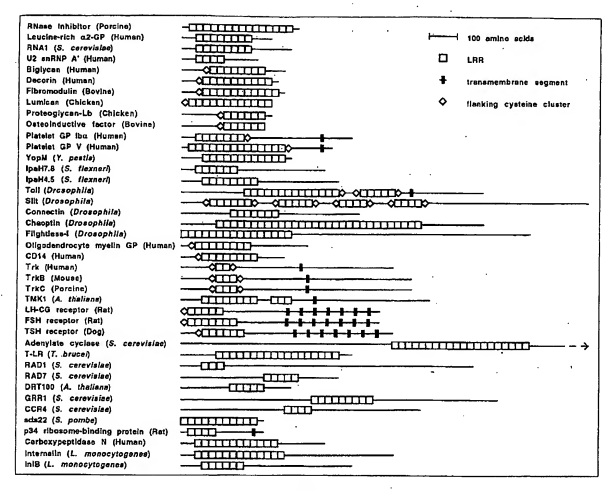


Figure 1
Schematic diagrams of the primary structures of proteins with leucine-rich repeats. For abbreviations and references, see Table I.

The consensus residues of the repeats play structural roles. The sidechains of leucines and of other aliphatic residues (positions 2, 5, 7, 12, 17, 20, 23 and 24), of cysteines (position 17), and of asparagines or cysteines (position 10) form the core of the protein [note that the consensus positions specific to RI are slightly different from the overall consensus of LRR-containing proteins (Table 1)]. The spacing of leucines is responsible for proper packing of the β -strands and α -helices. The remaining consensus residues in RI have other roles. For example, in the B-type repeat, glycine at position 13 of the repeat seems to be a helix-initiation signal, while proline at position 27 is a helix-termination signal. Because of the arrangement of the β-α units, sidechains of consensus residues align in stacks, both at the core and on the surface of the protein.

It is unlikely that a single leucine-rich motif would adopt the conformation observed in the RI protein. Neighboring repeats form many polar and nonpolar contacts; also, the β -strands tend to pack against the helices that precede and follow the β -strand in question. Synthetic peptides that correspond to a single repeat have low propensity to adopt a defined structure^{5,6}.

Secondary-structure predictions of LRRs show no clearly repetitive patterns. Consequently, before the structure of RI had been determined, both an α -helix, and a β -strand conformation, was proposed as the major secondary structural element of LRRs in different proteins. β -turns were, however, correctly predicted in the vicinity of the conserved asparagine.

Leucine-rich repeats should not be confused with the more widely known

leucine zippers. Leucine zippers are motifs found in oligomeric proteins, including many DNA-binding proteins such as the products of the c-fos and c-jun proto-oncogenes. Leucine zippers are built of heptad repeats with leucines present at every seventh position. These leucines form a hydrophobic ridge on one side of an a-helix that interacts with a complementary helix10. The only similarity between LRRs and leucine zippers is the conservation of leucines at particular spacings; these leucines cannot usually be replaced by other hydrophobic amino acids. However, the spacing between leucines differs between LRRs and leucine zippers, with only certain leucines in the carboxy-terminal part of LRRs spaced by seven residues. The function of leucines in LRRs and leucine zippers is also different; while leucines in leucine zippers participate in oligomerization, the buried leucines in

(a) (b) Α 8 G DxGaxxxxxxLxxPxx αβε Ben αα (c),

Figure 2

Structure of ribonuclease inhibitor (RI). (a) Ribbon diagram of the structure of porcine RI. (b) Consensus sequences and secondary structure of the A- and B-type leucine-rich repeats (LRRs) of porcine RI. One-letter amino acid code is used; 'x' indicates any amino acid, 'a' denotes an aliphatic amino acid. The part of the repeat that is strongly conserved throughout the LRR superfamily is underlined, and the conserved residues are bold. Below the sequence, elements of secondary structure are indicated. The solid line marks the core region of the β -sheet or α -hellx; the dots denote extensions of the helix in different repeats. The types of β -turns are shown below the corresponding i+1 and i+2 positions. (c) Stereo plot of the main chain of a representative leucine-rich repeat corresponding to residues 195–222 of porcine RI. Sidechains of only the consensus residues are shown; bonds in the conserved residues are filled.

LRRs play a structural role and cannot be directly involved in protein-protein interactions.

RI as a model for other LRR-containing proteins

We believe that the repeats in RI are a satisfactory structural model for LRRs in other proteins. Most consensus residues that play key structural roles are conserved throughout the superfamily, especially the consensus residues that cluster around the β -strand region (Table I, Fig. 2). Although the LRRs in RI are among the longest found in the superfamily, shorter repeats could easily adopt a similar β - α conformation, as demonstrated by the amino-terminal structural repeat in RI, which contains 25 residues.

We must, however, allow the possibility that in certain LRR-containing proteins the helical parts might be substituted with more extended structures. This would most probably occur in those proteins that contain shorter variants of LRRs which, in the carboxy-terminal regions of the repeats, lack the appropriate pattern of hydrophobic residues expected for an amphipathic helix, and contain prolines and other amino acids with low propensities to form α -helices.

Substitution of the helical part of the repeats with a more extended structure would influence the overall shape of the protein considerably, particularly its curvature. Two protein folds exist that could shed light on the plausible architecture of a structure with β -strands substituting for a-helix. These are the β-roll folds of pectate lyase C11 (PelC) and alkaline protease¹² (AP) (Fig. 3). Both β-roll folds are assembled from repetitive structural units and expose a surface of a parallel β -sheet to solvent. In addition, in PelC, as in RI, a ladder of asparagine residues at the carboxyterminal ends of the β -strands hydrogen bonds to the backbone. The length of a structural repeat of the PelC-type 'Bhelix' is 22 residues, and the length of a tum of the AP-type β -roll is 18 residues. Neither PelC nor AP contain LRRs.

In proteins that contain over 21 LRRs and retain the curvature of the structure of RI, the overall organization of the repeats must be different from that in RI to prevent the termini from colliding. Such a collision could be avoided by a twist in the sheet. Alternatively, smaller numbers of repeats could constitute separate domains, linked by segments adopting conformations different from the β - α units.

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LRR-containing proteins: functional and evolutionary similarities

LRRs are found in a functionally and evolutionarily diverse set of proteins. Although few properties are shared among all members of the LRR superfamily, two main themes emerge: all LRR-containing proteins appear to be involved in protein-protein interactions, and at least half of them take part in signal transduction pathways (Table I). LRRs must therefore provide a structural skeleton for the achievement of diverse. specific molecular interactions.

A property of the structure of the LRR that may be helpful in protein binding is its nonglobular shape. Because of this shape, the area available for interaction with smaller globular proteins is substantially increased, facilitating more interactions, and thus enhancing affinity.

Another property of the structure of LRRs that might be useful for protein binding is the exposed surface of the parallel β-sheet. This structural feature is rare in proteins, and has only been discovered recently in structures of PelC11 AP12 and RI2. The β-roll structures of PelC and AP were proposed to be responsible for transport across membranes. However, such a surface feature could also aid in protein binding. Parallel β -sheets are common in the interior of proteins, where helices or other parallel \beta-sheets pack against them. In the case of LRR-containing proteins, the interaction with the ligand could substitute for such packing interactions.

In addition to the properties common to most LRR-containing proteins, such as involvement in protein-protein interactions and signal transduction, further similarities restricted to fewer members of the superfamily can be found, on the basis of both their functions and the characteristics of their repeats. These often point to evolutionary and functional relationships among LRRcontaining proteins. The specificity of protein-protein interactions of LRR-containing proteins and, therefore, the basis of their functions are most probably due to the specific composition of nonconsensus residues, and might also be influenced by the lengths of the repeats and the flanking domains. These functional and evolutionary relationships are discussed below, and the remarkable diversity of functions of LRR-containing proteins is demonstrated.

Relatives of Ri. The closest known relatives of RI seem to be the RNA1 protein, which is involved in RNA processing 13,

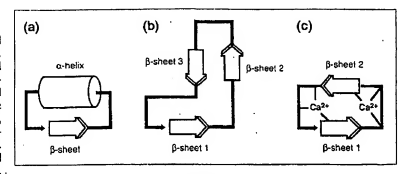


Figure 3 Schematic diagrams of the repetitive structural units of (a) ribonuclease inhibitor, (b) pectate lyase C, and (c) the β-roll domain of alkaline protease.

and the A' protein from the U2 small nuclear ribonucleoprotein particle (snRNP), which is required for snRNP assembly8. RI and RNAI also have the most closely related patterns of conserved residues among the LRR proteins. RI and A' protein both interact with RNA-binding proteins, and therefore may be functional relatives.

Adhesive proteins represent the largest group in the LRR superfamily. These proteins all contain similar 24-residue LRRs. One family of adhesive LRR-containing proteins includes the small proteoglycans: biglycan, fibromodulin, decorin, lumican, proteoglycan-Lb and osteoinductive factor (OIF, also called osteoglycin)9. Small proteoglycans bind various components of the extracellular matrix and growth factors (Table 1). Decorin and fibromodulin regulate collagen-fibril formation9; and OIF, in conjunction with the transforming growth factors TGF-β and TGF-β2, induces bone formation14.

Several adhesive LRR-containing proteins are found in Drosophila. Toll, slit, connectin, chaoptin and flightless-I appear to orient cells during development; each protein is specific to particular tissues 15-19. Connectin, chaoptin and a similar mammalian protein that may be involved in myelination (oligodendrocyte-myelin glycoprotein, OMGP)20 are attached to the membrane via a glycosylphosphatidylinositol (GPI) anchor.

Another family of adhesive proteins comprises the proteins of the Ib-V-IX system of platelet glycoproteins. This complex constitutes the receptor for von Willebrand factor (vWF) and mediates the adhesion of platelets to injured vascular surfaces21. All polypeptides in the complex contain LRRs, but glycoproteins IbB and IX contain only one leucine-rich motif. Although it is unlikely

that a single motif by itself would normally adopt the conformation observed in the crystal structure of RI, such a conformation could be stabilized by other proteins in the complex. Bacterial proteins YopM and IpaH interfere with platelet-mediated inflammatory responses, supposedly owing to homology with glycoprotein $lb\alpha^{22}$.

Signal-transducing receptors. The LRR superfamily contains several families of signal-transducing receptors. CD14 is a GPI-anchored receptor that, upon binding the complex between lipopolysaccharide and its binding protein, triggers intracellular protein tyrosine phosphorylation, which in turn leads to an antibacterial response by macrophages23.

The proto-oncogene trk encodes a transmembrane protein, Trk, with a cytoplasmic tyrosine kinase domain24. The extracellular domain that contains LRRs is involved in high-affinity binding of nerve growth factor; other proteins from the Trk family selectively bind other neurotrophins (Table I). TMK1, a plant protein with structural and biochemical properties similar to the receptor protein kinases, contains serine/ threonine kinase in its cytoplasmic domain²⁵. The receptor protein kinases are related to adhesive proteins such as Toll and glycoprotein lb, as they are targeted to the plasma membrane and contain an extracellular LRR domain, a transmembrane domain and a carboxyterminal cytoplasmic domain involved in signal transduction.

G-protein-coupled gonadotrophin receptors consist of an extracellular, hormone-binding LRR domain, a domain containing seven membrane-spanning segments and a short cytoplasmic tail26. Their ligands, chorionic gonadotrophin, luteinizing hormone, follicle-stimulating hormone and thyroid-stimulating

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hormone, are heterodimeric glycoproteins composed of a specific β -subunit noncovalently attached to a common α -subunit.

Other signal-transducing proteins. Yeast adenylate cyclase contains LRRs in a domain that is important for the regulation of its catalytic activity by the RAS protein²⁷. The trypanosomal leucine-rich protein may be related to this domain because it is encoded in the variant-surface-glycoprotein expression site adjacent to an adenylate cyclase²⁸.

Two LRR-containing proteins take part in the glucose repression pathway of yeast. GRR1 is the primary response element in this pathway²⁹, and CCR4 is required for the transcription of glucose-repressible alcohol dehydrogenase²⁰.

Other proteins. RAD1 and RAD7 constitute another functional group of LRR-containing proteins, as they are both involved in nucleotide excision repair in yeast. RAD1 forms a functional complex with RAD10, but the binding region lies outside the LRR-containing sequence³¹. Genetic evidence suggests that RAD7 may interact with RAD23 (Ref. 32).

RAD1 also plays a role in mitotic recombination³³. Similarly, DRT100, a plant cDNA encoding a protein that contains LRRs and a nucleotide-binding motif, complemented deficiencies in homologous recombination in Escherichia coli³⁴.

The remaining LRR-containing proteins thus far appear unrelated to other members of the superfamily, but also have important functions: sds22 regulates protein phosphatase-1 function in fission-yeast mitosis³⁵; the ribosome-binding protein p34 may function as the ribosome receptor in the secretory pathway³⁶; the two LRR-containing subunits of the carboxypeptidase N tetramer stabilize the catalytic subunits and keep them in circulation³⁷; and internalin confers invasive properties on Listeria³⁸.

Flanking cysteine clusters. Many LRR-containing proteins also contain homologous regions flanking the LRR domains (Fig. 1). These regions are characterized by four similarly spaced cysteines in a stretch of about 20 amino acids for the amino-flanking region and about 50 amino acids for the carboxy-flanking region 16.39; the consensus sequences can be described as CP[-2x]CxC[-6x]C for the amino-flanking and PxxCxC [-20x]C[-20x]C for the carboxy-flanking regions (x denotes any amino acid). The amino-flanking cysteine

clusters occur in all small proteoglycans, glycoproteins IbB and IX, Toll, slit, OMGP, CD14, the Trk family and gonadotrophin receptors; related sequences without all four cysteine residues conserved are present in leucine-rich a2-glycoprotein, glycoproteins $\mbox{Ib}\alpha$ and $\mbox{V, IpaH4.5, chaoptin}$ and carboxypeptidase N. The carboxyflanking cysteine clusters occur in glycoproteins $Ib\alpha$, $Ib\beta$, V and IX, Toll, slit and the Trk family; related sequences in which all four cysteines are not conserved are present in leucine-rich a2glycoprotein and connectin. A different type of carboxy-flanking sequence containing two cysteines is found in small proteoglycans39. The flanking cysteine clusters appear to be a property of adhesive proteins and receptors.

LRRs with known function. In several cases, LRRs are responsible for specific functions. In RI, RNA1, small proteoglycans, chaoptin, CD14 and sds22, the tandem repeats comprise essentially the entire primary structure. In glycoprotein lbα, the binding sites for vWF and thrombin have been mapped to the LRR-containing domain; although LRRs do not appear to bind vWF directly21, point mutations in the LRR domain cause variants of Bernard-Soulier disease, a bleeding disorder (see, for example, Ref. 40). LRRs appear to be responsible for the regulation of yeast adenylate cyclase by RAS27, for the hormone selectivity of the gonadotrophin receptors41 and for the function of CCR4 (Ref. 30). The single leucine-rich motif in the type 1 human immunodeficiency virus Rev protein is the trans-activating region of the virus⁴².

Concluding remarks

LRR-containing proteins represent an important superfamily involved in protein-protein interactions and signal transduction. The wide range of functions of LRR-containing proteins suggests that LRRs are generally useful proteinbinding motifs; the variability of primary structures of these proteins suggests that these proteins do not necessarily share a common ancestor. The crystal structure of RI provides a model for the architecture of LRRs and helps us understand why LRRs may be useful for protein binding; the two structural features that might be responsible for the binding function are the solvent-exposed parallel B-sheet and the nonglobular shape of the structure. Owing to the variability of repeats and functional diversities of LRR-containing proteins, however, the

crystal structure of RI can only represent a platform for further structural studies of these proteins.

Acknowledgements

We thank D. Crawford and B. Beutler for making us interested in ribonuclease inhibitor and leucine-rich repeats, and K. Fischer Lindahl and J. F. Sambrook for review of the manuscript and suggestions.

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CELLS ARE COMPOSED primarily of water, and movement of water into and

out of cells is a key feature of numerous .

physiological functions in animals, plants and microorganisms. The plasma mem-

branes of renal proximal tubules, certain other mammalian epithelia and erythro-

cytes are highly permeable to water, which moves in the direction of an

osmotic gradient (see Refs 1, 2 for reviews). In plants, water moves through the living tissues in response to hydro-

static and osmotic forces. During the day, when the stomata are open, a constant transpiration stream moves

through the tissues; water enters the

roots and moves up the conductive tis-

sues of the stem, and water vapor exits

meability of mammalian epithelia and

the hydraulic conductance of plant tissues and cells have been made, and the

plasma membranes are thought to be

the major barriers to water flow. The

lipid bilayer is somewhat permeable to water resulting from simple diffusion, and the diffusional water permeability

coefficient (P_d) of many membranes is

<0.01 cm s⁻¹. This process cannot be in-

hibited pharmacologically and is charac-

terized by a high Arrhenius activation

energy ($E_a > 10$ kcal mol⁻¹, i.e. reduced

water movement at lower temperatures

where lipid packing is tighter). Pores or

water channels were hypothesized to

explain the more rapid transmembrane

water movements characteristic of eryth-

rocytes, certain mammalian epithelia and plant-tissues. Unlike diffusional water

permeability, channel-mediated water

movement across erythrocyte mem-

branes is reversibly inhibited by mercuric

chloride and exhibits a low Arrhenius

activation energy ($E_a < 5 \text{ kcal mol}^{-3}$).

The osmotic water permeability

M. J. Chrispeels is at the Department of

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Numerous studies on the water per-

the leaves.

Proc. Natl Acad. Sci. USA 89, 8073-8077 35 Stone, E. M., Yamano, H., Kinoshita, N. and Yanagida, M. (1993) Curr, Biol, 3, 13-26

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Aquaporins: water channel proteins of plant and animal cells

Maarten J. Chrispeels and Peter Agre

Certain biological membranes, such as the erythrocyte plasma membrane, have a high osmotic water permeability, and such membranes have long been suspected of harboring water-channels. The molecular identity of these channels has now been established with the purification of water-channel proteins and the cloning of the genes encoding them. Homologous water-channel proteins, called 'aquaporins', are present in plants and animals. These channels are water selective and do not allow ions or metabolites to pass through them. Their discovery is providing new insights into how plant and animal cells facilitate and regulate the passage of water through their membranes.

coefficient (P,) of erythrocyte membranes measures approximately 0.02 cm sec-1. Although erythrocyte membranes and certain other biological membranes have $P_i >> P_d$, suggesting the existence of water channels, the molecular identity of the water channels remained elusive until the recent identification of CHIP and a growing list of evolutionarily related water-channel proteins in animals and plants.

An ancient family of channel proteins

The aquaporins are a functionally defined group of water-transport proteins that are part of an ancient family of channel proteins called the MIP family, after the first identified member, major intrinsic protein of the mammalian lens. MIP comprises approximately 50% of the total membrane protein of lens-fiber cells, and the cDNA encoding it was isolated from a bovine lens library4. Hydropathy analysis of the deduced amino acid sequence predicted a 263-residue integral membrane protein with cytoplasmic amino and carboxyl

termini and six bilayer-spanning domains. The amino- and carboxyterminal halves of MIP are themselves sequence related, forming a unique structure in which the topology of the first and second halves of the molecule are oriented at 180° to each other (Fig. 1); the MIP family already has more than 20 members, including: NOD26, a protein in the symbiosome membrane of soybean root nodules; the glycerol facilitator of Escherichia coli and other bacteria; a suppressor of a Saccharomyces cerevisiae growth defect on fermentable sugars; a developmentally regulated Drosophila brain protein; and multiple tonoplast intrinsic proteins in plants (Ref. 5 and references therein). The functions of many MIP family members have yet to be established, but several plant and animal MIP-family members have been shown to be water-specific membrane channels.

Structure of the aquaporins

As with MIP, the 28kDa channelforming integral protein (CHIP) was first

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